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/050182 A

(54) Title: ELASTOMER COMPOSITES, ELASTOMER BLENDS AND METHODS

(57) Abstract: Elastomer composites and elastomer blends comprising particulate fillers having selected and surface area values are disclosed as well as elastomer composites and elastomer blends having improved properties. Methods of making and using elastomer composites and elastomer blends are also disclosed.

#### **ELASTOMER COMPOSITES, ELASTOMER BLENDS AND METHODS**

#### **Field of Invention**

[0001] Certain aspects of the present invention relate to elastomer composites and elastomer blends. Other aspects relate to novel methods of making elastomer composites and elastomer blends.

#### Background

Numerous products of commercial significance are formed from elastomeric compositions wherein particulate filler is dispersed in various synthetic elastomers, natural rubber or elastomer blends. Carbon black, for example, is widely used as a reinforcing agent in natural rubber and other elastomers. Certain grades of commercially available carbon black are used, which vary both in surface area per unit weight and in structure, but have been limited by conventional rubber batch processing techniques. Numerous products of commercial significance are formed from such elastomeric compositions, including, for example, vehicle tires, engine mount bushings, conveyor belts, windshield wipers and the like. While a wide range of performance characteristics can be achieved employing currently available materials and manufacturing techniques, there has been a long standing need in the industry to develop elastomeric compositions having improved properties, especially elastomer compositions that can be produced effectively and economically.

#### **Summary**

[0003] In accordance with a first aspect of the invention, there is provided an elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer. The particulate filler comprises at least one carbon black having structure and surface area values which satisfy the equation CDBP  $\leq$  (BET  $\div$  2.9)-X, wherein X is greater than or equal to 0. Such carbon blacks are referred to in some instances below as ultra-high surface area/low structure carbon blacks. Preferably, the

elastomer is natural rubber. Additional ingredients, e.g., any of numerous additives and other fillers known for use in elastomer composites, may be included in the elastomer composites, such as to achieve desired performance properties, processing characteristics etc.

[0004] In accordance with another aspect of the invention, there is provided an elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer having a tear strength, as measured for example, by test method ASTM-D624 using Die C of greater than about 160 N/mm, more preferably greater than about 165 N/mm, and most preferably greater than about 170 N/mm.

[0005] In accordance with another aspect of the invention, there is provided an elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer having a Shore A hardness, as measured for example in accordance with test method ASTM-D1415, of greater than about 65, a tensile strength, as measured for example in accordance with test method ASTM-D412, of greater than about 30 megapascals, and an elongation at break, as measured for example in accordance with test method ASTM-D412 of greater than about 600 %.

[0006] In accordance with another aspect of the invention, there is provided an elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer, wherein the particulate filler comprises at least one carbon black in an amount such that the elastomer has tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm.

[0007] In accordance with another aspect of the invention, there is provided an elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer, wherein the particulate filler comprises at least one carbon black in an amount such that the elastomer has Shore A hardness greater than about 65, as measured in accordance with test method ASTM-D1415, tensile strength greater than about 30 megapascals, as measured in accordance with test method ASTM-D412, and elongation at break greater than about 600 %, as measured in accordance with test method ASTM-D412.

[0008] In accordance with another aspect of the invention, there is provided an elastomer composite comprising particulate filler dispersed in elastomer by methods referred to here as continuous wet mixing and coagulation wherein the

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carbon black has structure and surface area values satisfying the equation CDBP ≤ (BET ÷ 2.9)-X, wherein X is greater than or equal to 0. Certain preferred embodiments of such elastomer composites have tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm. Certain other preferred embodiments of such elastomer composites have Shore A hardness, as measured in accordance with test method ASTM-D1415, greater than about 65, tensile strength, as measured in accordance with test method ASTM-D412, greater than about 30 megapascals, and elongation at break, as measured in accordance with test method ASTM-D412, greater than about 600 %.

[0009] In accordance with another aspect of the invention, there is provided elastomer blends comprising at least one elastomer composite disclosed above blended with an elastomer material. The elastomer material comprises at least one elastomer, which may be the same as, or different from, the elastomer in the elastomer composite. Optionally, the elastomer material comprises filler, e.g., carbon black, additives or other fillers dispersed in the elastomer.

In accordance with yet another aspect of the invention, there is provided a method of preparing the elastomer composites and elastomer blends disclosed above. This method comprises compounding or mixing the elastomer and the particulate filler, including the carbon black, and optionally other ingredients. In certain preferred embodiments such elastomer composites are produced by methods comprising continuous wet mixing and coagulation. Similarly, in certain preferred embodiments such elastomer blends are produced by methods comprising continuous wet mixing and coagulation followed by further compounding or mixing with additional elastomer material comprising at least one elastomer, which may the same as, or different from, the elastomer in the elastomer composite. Optionally, the elastomer material comprises filler, e.g., carbon black, additives or other fillers dispersed in the elastomer.

[0011] These and other aspects and advantages will be further understood in view of the following detailed discussion of certain preferred embodiments.

#### **Detailed Description of Certain Preferred Embodiments**

Preferred methods and apparatus for producing elastomer composites [0012] disclosed here are described in commonly assigned United States Patent Nos. 6,075,084, 6,048,923, and 6,040,364, the entire disclosure of each of which is hereby incorporated herein by reference for all purposes. Such methods are referred to here in some instances as continuous wet mixing and coagulation. A preferred method for producing the elastomer composites comprises feeding a continuous flow of first fluid comprising elastomer latex to a mixing zone of a coagulum reactor defining an elongate coagulum zone extending from the mixing zone to a discharge end, and feeding a continuous flow of second fluid comprising particulate filler under pressure to the mixing zone of the coagulum reactor to form a mixture with the elastomer latex. The mixture passes as a continuous flow to the discharge end of the coagulum reactor, and the particulate filler is effective to coagulate the elastomer latex. More specifically, the second fluid is fed against the first fluid within the mixing zone sufficiently energetically to substantially completely coagulate the elastomer latex with the particulate filler prior to the discharge end of the coagulum reactor. A substantially continuous flow of elastomer composite is discharged from the discharge end. As noted above, these processes are referred to below in some instances as continuous wet mixing and coagulation.

In certain preferred embodiments, particulate filler slurry is fed to the mixing zone preferably as a continuous, high velocity jet of injected fluid, while the latex fluid typically is fed at relatively lower velocity. The high velocity, flow rate and particulate concentration of the filler slurry are sufficient to cause mixture and high shear of the latex fluid, flow turbulence of the mixture within at least an upstream portion of the coagulum zone, and substantially complete coagulation of the elastomer latex prior to the discharge end. Substantially complete coagulation can thus be achieved, in accordance with preferred embodiments, without the need of employing an acid or salt coagulation agent. Preferred continuous flow methods for producing the elastomer composites comprise continuous and simultaneous feeding of latex fluid and filler slurry to the mixing zone of the coagulum reactor, establishing a continuous, semi-confined flow of a mixture of the latex and filler slurry in the coagulum zone. Elastomer composite crumb in the form of "worms" or globules are

discharged from the discharge end of the coagulum reactor as a substantially constant flow concurrently with the on-going feeding of the latex and carbon black slurry streams into the mixing zone of the coagulum reactor. Feed rates of the latex fluid and carbon black slurry to the mixing zone of the coagulum reactor can be precisely metered to achieve high yield rates, with little free latex and little undispersed carbon black in the product crumb at the discharge end of the coagulum reactor. High feed velocity of the carbon black slurry into the mixing zone of the coagulum reactor and velocity differential relative the latex fluid feed are believed to be significant in achieving sufficiently energetic shear of the latex by the impact of the particulate filler fluid jet for thorough mixing and dispersion of the particulate into the latex fluid and coagulation of the latex. Prior techniques involving premixing of latex and particulate filler, such as in the above-mentioned Heller et al patent and Hagopian et al patent, do not recognize the possibility of achieving coagulation without exposing the latex/particulate mixture to the usual coagulant solution with its attendant cost and waste disposal disadvantages. Modified and alternative suitable methods for producing novel elastomer composites disclosed here will be apparent to those skilled in the art, given the benefit of this disclosure.

[0014] Numerous carbon blacks are suitable for use in the elastomer composites disclosed here, including commercially available carbon blacks and fillers comprising carbon black. In addition to the carbon blacks specifically disclosed here, additional carbon blacks will be apparent to those skilled in the art, given the benefit of this disclosure.

[0015] In accordance with one aspect of this invention, there is provided an elastomer composite comprising carbon blacks having surface area and structure values satisfying equation (1):

$$CDBP \le (BET \div 2.9) - X \tag{1}$$

wherein X is greater than or equal to 0. Preferably X is 0, and, when X is 0, Equation (1) may also be represented herein as "CDBP  $\leq$  (BET  $\div$  2.9)."

[0016] In Equation (1), the structure value CDBP is the dibutylphthalate adsorption number after the sample has been crushed and is measured in accordance with the test procedure described in ASTM D-3493. The surface area value BET is nitrogen adsorption surface area and is measured in accordance with the test

procedure described in ASTM D-4820. An example of a carbon black which satisfies this equation is BP1100, which has a CDBP value of about 43 to 45 mL/100g and a BET value of about 260 to 264 m²/g. Other preferred carbon blacks meeting Equation (1) include, for example, BP 1180, BP880, and CSX439, It has now been found that certain preferred embodiments of elastomer composites disclosed here, comprising such ultra-high surface area/low structure carbon blacks, have advantageous performance properties and processing characteristics.

[0017] In accordance with certain preferred embodiments, elastomer composites are disclosed here comprising carbon blacks having surface area and structure values satisfying Equation (2):

$$CDBP \le (BET \div 2.9) - X \tag{2}$$

wherein X is preferably about 5. More preferred are carbon blacks having surface area and structure values satisfying Equation (2) wherein X is about 10. It will be recognized, that the carbon blacks which satisfy Equation (2) are a subset of the carbon blacks which satisfy Equation (1). Additional suitable ultra-high surface area/low structure will be apparent to those skilled in the art, given the benefit of this disclosure.

[0018] Preferably, there is at least about 60 phr of ultra-high surface area/low structure carbon black in the elastomer composite. More preferably, there is at least about 65 phr of such ultra-high surface area/low structure carbon black in the elastomer composite, e.g., at least about 70 phr of such ultra-high surface area/low structure carbon black. Particularly preferred are such elastomer composites prepared by continuous wet mixing and coagulation and elastomer blends prepared by continuous wet mixing and coagulation and follow-on dry mixing with additional elastomer and/or other fillers, additives, etc. The advantageous elastomer composite properties disclosed here, such as high tensile strength, e.g., tensile strength over 160 N/mm, and good hardness, tensile strength and elongation, for example, are not found using traditional reinforcing carbon blacks.

[0019] The tear strength of the natural rubber elastomer composites comprising ultra-high surface area/low structure carbon black, e.g., carbon black BP1100, prepared by continuous wet mixing and coagulation exceeds the tear strength, as measured by test method ASTM-D624 using Die C, of natural rubber

elastomer composites of the same formulation but produced by dry mixing. At least certain preferred embodiments of continuous wet mixing and coagulation elastomer composite comprising BP 1100 have now been found to achieve a value of 160 N/mm at less than about 65 phr filler and to exceed 160 N/mm at higher loading levels. The carbon black BP 1100 is commercially available from Cabot Corporation and, as noted above, has surface area and structure values meeting the equation CDBP  $\leq$  (BET  $\div$  2.9). More specifically, as described above, BP 1100 has a BET surface area value of about 260 to 264 m²/g and a CDBP structure value of 43 to 45 mL/100g. Also, it has now been found that the tear strength of such a BP 1100 continuous wet mixing and coagulation elastomer composite significantly exceeds the tear strengths of comparably formulated elastomer composites produced by dry-mixing with other commercially available carbon blacks not meeting the equation CDBP  $\leq$  (BET  $\div$  2.9).

[0020] Comparative tear strength data also shows that the tear strength of elastomer composites comprising ultra-high surface area/low structure carbon black BP 1100, and produced by continuous wet mixing and coagulation, exceeds the tear strength of comparable elastomer composite comprising carbon black V7H, a non-ultra-high surface area/low structure carbon black, and produced by dry-mixing. Comparative tear strength data also shows that the tear strength of elastomer composites comprising BP 1100 and produced by continuous wet mixing and coagulation exceeds, at those tested loading levels between 50 and 110 phr, the tear strength of comparable elastomer composites comprising BP 1100 but produced by dry-mixing.

[0021] At least certain elastomer composites in accordance with this disclosure are suitable to be blended with additional elastomer, filler, other additives, etc. That is, at least certain of the elastomer composites disclosed here can be blended by subsequent dry-mixing with additional elastomer and/or filler or other additives, etc., including additional elastomer composites of the present invention Elastomer materials blended with elastomer composites disclosed here optionally comprise the same or different elastomer, and optionally may have carbon black and/or other filler or other additives dispersed therein.

[0022] Elastomer composites of the invention, particularly those prepared by continuous wet mixing and coagulation, can be incorporated into elastomer blends

using a variety of techniques known in the art, including by subsequent dry mixing with additional elastomer material, e.g. a second elastomer or an elastomer composite already comprising fillers and/or additives, etc. The dry mixing can be carried out with any suitable apparatus and techniques, such as commercially available apparatus and techniques. In one embodiment, aBanbury mixer or the like is used. Other ingredients also may be added along with the additional elastomer during dry mixing, including, for example, extender oil, antioxidant, cure activators, additional particulate filler, curatives (for example, zinc oxide and stearic acid), etc. In those embodiments wherein additional filler is added during follow-on dry mixing, such additional filler can be the same as or different from filler(s) in the elastomer composite. The elastomer blends may optionally undergo further processing steps known to those skilled in the art.

Preferred methods of producing elastomer composite blend, [0023] comprising first preparing elastomer composite by continuous wet mixing and coagulation as described above, followed by dry mixing the elastomer composite with additional elastomer material to form elastomer blend, are described in U.S. Patent No. 6,075,084, the entire disclosure of which is hereby incorporated herein by reference for all purposes. This process may be referred to in some instances as continuous wet mixing and coagulation with follow-on dry mixing. As stated above, at least certain preferred embodiments of the elastomer composites disclosed here are produced by continuous wet mixing and coagulation, and at least certain preferred embodiments of elastomer blends can be produced by continuous wet mixing and coagulation with follow-on dry mixing. In accordance with certain preferred embodiments, elastomer composites are produced by such continuous wet mixing and coagulation methods and apparatus, in a continuous flow process without the need for using traditional coagulating agents, such as acids or salts.

[0024] Advantageous flexibility is achieved by the method disclosed here for making elastomer blends, comprising continuous wet mixing and coagulation with follow-on dry mixing. In particular, flexibility is provided as to the choice of elastomer(s) employed in continuous wet mixing and coagulation and in the choice of elastomer(s) used in the subsequent dry mixing step. The same elastomer or mixture of elastomers can be used in the wet and dry mixing steps or, alternatively, different

elastomers can be used in any suitable relative weight proportion. Further flexibility is provided in that additional filler and other additives and the like may optionally be added during either wet mixing or dry mixing. It should be understood that the dry mixing can be a multi-stage compounding process. Such additional materials can be the same as or different from those used in the continuous wet mixing and coagulation. Without wishing to be bound by theory, it presently is understood that, in at least certain preferred embodiments, a multi-phase elastomer composite blend is produced by continuous wet mixing and coagulation with follow-on dry mixing. That is, although difficult to identify or observe using techniques currently in general use in the elastomer industry, the elastomer blend is understood to comprise at lease one elastomer phase produced by continuous wet mixing and coagulation and another elastomer phase added or produced by follow-on dry mixing. The degree of mixing or blending of the two phases and the degree to which boundary layers between the two phases are more or less distinct will depend on numerous factors, including, for example, the mutual affinity of the elastomers, the level of filler loading, the choice of filler(s) and whether additional filler is added during dry mixing, the relative weight proportion of the continuous wet mixing and coagulation elastomer and the dry mixing elastomer, etc.

Numerous elastomers suitable for use in the elastomer composites [0025] disclosed here are commercially available or are otherwise known and prepared according to known techniques. Suitable elastomers include, but are not limited to, natural rubber, which is preferred, and other rubbers and polymers (e.g., homopolymers, copolymers, terpolymers, etc., all referred to here generally as polymers or copolymers unless otherwise stated or otherwise clear from context) of 1,3-butadiene, styrene, isoprene, isobutylene, 2,3-dimethyl-1,3-butadiene, acrylonitrile, ethylene, and propylene and the like. In accordance with certain preferred embodiments, the elastomer has a glass transition temperature (Tg), as measured by differential scanning calorimetry (DSC), ranging from about -120° C to about 0°C. Examples include, but are not limited to, natural rubber and its derivatives such as chlorinated rubber, styrene-butadiene rubber (SBR), polybutadiene, polyisoprene, poly(stryene-co-butadiene) and the oil extended derivatives of any of them. Blends of any of the foregoing may also be used.

[0026] In certain preferred embodiments employing continuous wet mixing and coagulation, suitable elastomers are employed as latex fluids, e.g., natural or synthetic elastomer latices and latex blends. The latex preferably is suitable for coagulation by the selected particulate filler and must be suitable for the intended purpose or application of the final rubber product. It will be within the ability of those skilled in the art to select suitable elastomer latex or a suitable blend of elastomer latices for use in continuous wet mixing and coagulation to produce elastomer composites disclosed here, given the benefit of this disclosure. Exemplary elastomers include, but are not limited to, natural rubber and latices of the other elastomers recited above. The latex may be in an aqueous carrier liquid. Alternatively, the liquid carrier may be a hydrocarbon solvent. In any event, the elastomer latex fluid must be suitable for controlled continuous feed at appropriate velocity, pressure and concentration into the mixing zone. Suitable synthetic rubber latices include, for example, copolymers of from about 10 to about 70 percent by weight of styrene and from about 90 to about 30 percent by weight of butadiene, such as copolymer of 19 parts styrene and 81 parts butadiene, a copolymer of 30 parts styrene and 70 parts butadiene, a copolymer of 43 parts styrene and 57 parts butadiene and a copolymer of 50 parts styrene and 50 parts butadiene; polymers and copolymers of conjugated dienes such as polybutadiene, polyisoprene, polychloroprene, and the like, and copolymers of such conjugated dienes with an ethylenic group-containing monomer copolymerizable therewith such as styrene, chlorostyrene, acrylonitrile, styrene. 2-vinyl-pyridine, vinylpyridine, 5-ethyl-2-vinylpyridine, 2-methyl-5-vinylpyridine, alkyl-substituted acrylates, vinyl ketone, methyl isopropenyl ketone, methyl vinyl either, and alphamethylene carboxylic acids and the esters and amides thereof, such as acrylic acid and dialkylacrylic acid amide. Also suitable are copolymers of ethylene and other high alpha olefins such as propylene, 1-butene and 1-pentene.

[0027] Elastomers suitable for use in elastomer blends disclosed here, that is, elastomers suitable for addition to the disclosed elastomer composites, include numerous elastomers that are commercially available or are otherwise known and prepared according to known techniques. Exemplary elastomers include those listed above for use in the elastomer composites. In certain preferred embodiments of the

elastomer blends, specifically, those produced by continuous wet mixing and coagulation and follow-on dry mixing with additional elastomer, the additional elastomer during the dry mixing step can be any elastomer or mixture of elastomers suitable to the intended use or application of the finished product, including those listed above for use in continuous wet mixing and coagulation. In accordance with certain preferred embodiments, the elastomer latex employed in continuous wet mixing and coagulation is natural rubber latex and the additional elastomer employed in the dry mixing step is butadiene rubber (BR). In such preferred embodiments, the butadiene rubber preferably forms the minor phase or constituent of the elastomer composite blend, most preferably being from 10% to 50% by weight of total elastomer in the elastomer composite blend. In accordance with certain other preferred embodiments, the elastomer latex employed in continuous wet mixing and coagulation is natural rubber latex and the additional elastomer employed in the dry mixing step is styrene-butadiene rubber (SBR). In such preferred embodiments, the SBR preferably forms the major phase or constituent of the elastomer composite blend, most preferably being from 50% to 90% by weight of total elastomer in the elastomer composite blend. In accordance with certain other preferred embodiments, the additional elastomer is natural rubber. In accordance with certain other preferred embodiments, the elastomer latex employed in continuous wet mixing and coagulation is butadiene rubber latex and the additional elastomer employed in the dry mixing step is SBR. In such preferred embodiments, the SBR preferably is from 10% to 90% by weight of the total elastomer in the elastomer composite blend. In accordance with certain other preferred embodiments, the elastomer latex employed in continuous wet mixing and coagulation is butadiene rubber latex and the additional elastomer employed in the dry mixing step is natural rubber. In such preferred embodiments, the natural rubber preferably is the minor constituent or phase of the elastomer composite blend, most preferably being from 10% to 50% by weight of total elastomer in the elastomer composite blend. In accordance with certain other preferred embodiments employing butadiene rubber latex in continuous wet mixing and coagulation, the additional elastomer is additional butadiene rubber. In accordance with certain other preferred embodiments, the elastomer latex employed in continuous wet mixing and coagulation is SBR and the additional elastomer is

butadiene rubber. In such preferred embodiments, the butadiene rubber preferably is from 10% to 90% by weight of total elastomer in the elastomer composite blend. In accordance with certain other preferred embodiments, the elastomer latex employed in continuous wet mixing and coagulation is SBR and the additional elastomer is natural rubber. In such preferred embodiments, the natural rubber preferably is the major constituent or phase, most preferably being from 50% to 90% by weight of total elastomer in the elastomer composite blend. Certain other preferred embodiments SBR is employed in both the wet mixing and dry mixing steps, thus being essentially 100% of the elastomer in the elastomer composite blend.

[0028] Where the elastomer latex employed in continuous wet mixing and coagulation comprises natural rubber latex, the natural rubber latex can comprise field latex or latex concentrate (produced, for example, by evaporation, centrifugation or creaming). The natural rubber latex in such embodiments preferably is suitable for coagulation by the carbon black. The latex is provided typically in an aqueous carrier liquid. Alternatively, the liquid carrier may be a hydrocarbon solvent. In any event, the natural rubber latex fluid must be suitable for controlled continuous feed at appropriate velocity, pressure and concentration into the mixing zone. The wellknown instability of natural rubber latex is advantageously accommodated in certain preferred embodiments employing continuous wet mixing and coagulation, wherein it is subjected to relatively low pressure and low shear throughout the system until it is entrained into a semi-confined turbulent flow upon encountering a carbon black slurry feed stream jet of high velocity and kinetic energy in the mixing zone of a coagulum reactor. In certain preferred embodiments, for example, the natural rubber is fed to the mixing zone at a pressure of about 5 psig, at a feed velocity in the range of about 3-12 ft. per second, more preferably about 4-6 ft. per second. Selection of a suitable latex or blend of latices will be well within the ability of those skilled in the art given the benefit of the present disclosure and the knowledge of selection criteria generally well recognized in the industry.

[0029] As disclosed above, certain preferred embodiments of the elastomer composites of the invention comprise carbon blacks capable of providing the elastomer composite advantageous tear strength properties. In accordance with certain preferred embodiments, elastomer composites disclosed here comprise general

purpose rubber and particulate filler dispersed in the elastomer, wherein the particulate filler comprises at least one carbon black effective or sufficient in selected concentrations in the elastomer to achieve tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm. Examples of a general purpose rubber include, but are not limited to, natural rubber, polyisoprene, polybutadiene, styrene-butadiene rubber, nitrile-butadiene rubber, or ethylene-propylene rubber (including EPDM). Preferably such general purpose rubber is natural rubber.

[0030] As disclosed above, certain preferred embodiments of the elastomer composites disclosed here comprise carbon blacks capable of providing the elastomer composite advantageous hardness, tensile strength and elongation at break. In accordance with certain preferred embodiments, elastomer composites disclosed here comprise elastomer and particulate filler dispersed in the elastomer, wherein the particulate filler comprises at least one carbon black effective or sufficient in selected concentrations in the elastomer to achieve:

Shore A hardness, measured in accordance with test method ASTM-D1415, greater than about 65;

tensile strength, measured in accordance with test method ASTM-D412, greater than 30 megapascals; and

elongation at break, measured in accordance with test method ASTM-D412, of at least about 600 %.

[0031] In certain such preferred embodiments, the elastomer composite comprises elastomer and particulate filler dispersed in the elastomer, and the elastomer composite has Shore A hardness greater than about 65, tensile strength greater than about 30 megapascals, and elongation at break greater than about 600 %. Preferred embodiments comprise at least one ultra-high surface area/low structure carbon black, more preferably at least one ultra-high surface area/low structure carbon black.

[0032] Optionally, the filler employed in the elastomer composite comprises one or more materials additional to the aforesaid carbon black. In embodiments prepared by continuous wet mixing and coagulation, and for embodiments of the elastomer blends disclosed here prepared by continuous wet mixing and coagulation and follow-on dry mixing, the carbon black filler of the elastomer composite can

include also other material that can be slurried and fed to the mixing zone in accordance with the principles disclosed here. Suitable additional materials include, for example, conductive fillers, reinforcing fillers, fillers comprising short fibers (typically having an L/D aspect ratio less than 40), flakes, etc. Thus, exemplary particulate fillers which can be employed in elastomer composites disclosed here include, for example, other carbon blacks, fumed silica, precipitated silica, coated carbon blacks such as silica-coated carbon blacks, modified carbon blacks such as those having attached organic groups, and treated carbon blacks including metaltreated carbon blacks (for example silicon-treated carbon blacks), either alone or in combination with each other. Suitable modified carbon blacks include those disclosed in U.S. Patent Nos. 5,851,280, 5,672,198, 6,042,643, 5,900,029, and 5,559,169 and U.S. Patent Application Serial No. 09/257,237, the entire disclosures of which are hereby incorporated by reference for all purposes. Suitable treated carbon blacks and coated carbon blacks are known and include those disclosed in U.S. Patent Nos. 5,916,934, 5,830,930, 6,028,137, 6,008,272, 5,919,841, 6,017,980, 5,904,762, 6,057,387, and 6,211,279 and U.S. Patent Application Serial Nos. 09/392,803 and 09/813,439, the entire disclosures of which are hereby incorporated by reference herein for all purposes. For example, in such silicon-treated carbon blacks, a silicon containing species, such as an oxide or carbide of silicon, is distributed through at least a portion of the carbon black aggregate as an intrinsic part of the carbon black. Also for example, in such silicon-coated carbon blacks, a silicon containing species, such as silica, is disposed on at least a portion of the surface of the carbon black aggregates. Additional materials and additives suitable to be employed with the aforesaid carbon blacks in the elastomer composites disclosed here will be apparent to those skilled in the art, given the benefit of this disclosure.

[0033] The rubber compositions of the present invention may optionally contain various additives along with the elastomer and filler, such as curing agents, coupling agents, and optionally, various processing aids, oil extenders and antidegradents. Examples of additives include, but are not limited to, antiozonants, antioxidants, plasticizers, resins, flame retardants, and lubricants. Combinations of additives can also be used. In that regard, it should be understood that the elastomer composites of the invention include vulcanized compositions (VR), thermoplastic

vulcanizates (TPV), thermoplastic elastomers (TPE) and thermoplastic polyolefins (TPO). TPV, TPE, and TPO materials are further classified by their ability to be extruded and molded several times without substantial loss of performance characteristics. Thus, in making or further processing the elastomer composite blends, one or more curing agents such as, for example, sulfur, sulfur donors, activators, accelerators, peroxides, and other systems used to effect vulcanization of the elastomer composition may be used.

[0034] The elastomer composite produced by continuous wet mixing and coagulation may optionally undergo further processing. For example, the elastomer composite may be further processed in a mixing and compounding apparatus, such as a continuous compounder. Suitable continuous compounders are described in PCT Publication No. WO 00/62990, the entire disclosure of which is hereby incorporated herein by reference for all purposes.

A significant advantage has now been recognized in preparing [0035] elastomer composites by continuous wet mixing and coagulation. excellent abrasion resistance is achieved, even in elastomer composites comprising oil to reduce hardness. Typically, hardness of an elastomer composite increases as the amount of filler in the composite is increased. Often, an intended use of an elastomer composite calls for high carbon black loading. However, the intended use of the elastomer composite may also call for lower hardness. It is known to add oil to reduce hardness of an elastomer composite, but if the amount of oil in an elastomer is increased to avoid undesirable hardness, the abrasion resistance of the elastomer composite generally is reduced. Elastomer composites prepared by continuous wet mixing and coagulation, including at least certain preferred embodiments of the disclosed here, achieve high abrasion resistance, elastomer composites notwithstanding high filler content and correspondingly high amounts of oil to control hardness. That is, abrasion resistance is found to be higher than in corresponding dry mixed elastomer composites. For example, novel elastomer composites made by continuous wet mixing and coagulation with loading levels of Vulcan 7H greater than 50 phr have higher abrasion resistance than corresponding dry mixed elastomer composite of the same formulation. It should be understood that the absolute value of abrasion resistance will depend on choice of filler, elastomer and oil, as well as filler

and oil loading levels, etc. For comparable formulations, however, elastomer composites produced by continuous wet mixing and coagulation, such as certain preferred embodiments of the elastomer composites disclosed here, have advantageously higher abrasion resistance than corresponding elastomer composites of the same formulation prepared using dry mixing techniques in accordance with the best commercial practices.

[0036] Elastomer composites were prepared using continuous wet mixing and coagulation and tested for abrasion resistance. Abrasion resistance was found generally to increases to a maximum and then decreases with increasing filler loading. It is also seen that the abrasion resistance of continuous wet mixing and coagulation elastomer composites is generally greater than that of corresponding drymixed elastomer composite, especially at the higher carbon black loading levels. For example, the abrasion resistance of a natural rubber elastomer composite prepared using continuous wet mixing and coagulation and carbon black BP1100 and H65 oil was found to be greater than the abrasion resistance of a comparable elastomer composite prepared using dry mixing methods. At 50 phr filler loading, the elastomer composite prepared using continuous wet mixing and coagulation exhibits a 200% increase in abrasion resistance over that of the analogous elastomer composite prepared using dry mixing. The abrasion resistance of natural rubber elastomer composites over a series of filler loading levels, prepared by continuous wet mixing and coagulation and employing carbon black Vulcan7H and H65 oil was found to be greater than the abrasion resistance of comparable elastomer composites prepared using dry mixing methods. The abrasion resistance of natural rubber elastomer composites over a series of filler loading levels, prepared by continuous wet mixing and coagulation and employing carbon black BP 1100, was found to be greater than the abrasion resistance of comparable elastomer composites prepared using dry mixing methods. At 80 phr filler loading, the elastomer composite prepared using the continuous wet mixing and coagulation exhibits 300% greater abrasion resistance than that of the corresponding elastomer composite prepared by dry mixing. Thus, using the continuous wet mixing and coagulation, especially employing ultra-high surface are/low structure carbon blacks, elastomer composites having high abrasion resistance can be produced.

[0037] As used here, the carbon black structure can be measured as the dibutyl phthalate adsorption (DBPA) value, expressed as cubic centimeters of DBPA per 100 grams carbon black, according to the procedure set forth in ASTM D2414. The carbon black surface area can be measured as CTAB expressed as square meters per gram of carbon black, according to the procedure set forth in ASTM D3765-85. Measurements of BET and CDBP values are as described previously above.

#### **EXAMPLES**

[0038] For each of the following examples, elastomer composite was produced comprising natural rubber from field latex, BP1100 carbon black available from Cabot Corporation, and aromatic oil. The properties of the natural rubber field latex are provided in Table 1 below:

Table 1
Natural rubber field latex properties

Total solids content, % (m/m)	32.8
Dry rubber content, % (m/m) (50:50 Ethanol/acetic acid)	31.9
Total alkalinity, NH3, % (g/100g latex)	0.510
VFA g KOH equiv. to the VFA in 100g latex solids	0.053
Acetone extract	2.11
Mooney viscosity ML (1+4)@100°C	90
Mw	1565976
Mn	1170073

[0039] The full formulation of the elastomer composite is set forth in Table 2 below.

Table 2 Formulations

Ingredient	phr
Rubber	100
BP1100	50-110
Aromatic oil	0-30
ZnO	4

Stearic Acid	2
6PPD (antioxidant)	1
TBBS (accelerator)	1.2
Sulfur	1.8

#### Examples 1-3

[0040] The following procedure, which is similar to that described in U.S. Patent No. 6,048,923, was used to prepare the elastomer composites of Examples 1-3.

#### 1. Carbon black slurry preparation

[0041] Bags of carbon black were dry ground and subsequently mixed with water in a carbon black slurry tank equipped with an agitator to form a 16.8 wt% carbon black slurry. This crude slurry was then fed to a homogenizer at an operating pressure of about 3000 psig such that the slurry was introduced as a jet into the mixing zone at a flow rate of about 780 kg/hr, to produce a finely ground carbon black slurry.

#### 2. Latex delivery

[0042] The latex, which was initially charged to a tank, was pumped to the mixing zone of the coagulum reactor. The latex flow rate was adjusted in order to obtain the desired final carbon black loading levels. Latex flow rates of between 430 to 600 kg/hr gave carbon black loading levels of between 80 and 95 phr (higher latex flow rates giving lower black loading levels). No antioxidant and oil were added in the latex.

#### 3. Carbon black and latex mixing

[0043] The carbon black slurry and latex were mixed by entraining the latex into the carbon black slurry. During entrainment, the carbon black was intimately mixed into the latex and the mixture coagulated. Soft, wet spongy "worms" of coagulum exited the coagulum reactor.

#### 4. Dewatering

[0044] The wet crumb discharged from the coagulum reactor was dewatered to 10 to 25 % moisture with a dewatering extruder (The French Oil Machinery Company, 7 inch diameter). In the extruder, the wet crumb was compressed and water squeezed from the crumb and through a slotted barrel of the extruder.

#### 5. Drying and cooling

[0045] The dewatered crumb was dropped into a continuous mixer where it was masticated and mixed with oil and antioxidant. Product exit temperature was less than 160°C and the moisture content was about 2%.

#### Tear Strength

[0046] Tear test results for the elastomer composites of Examples 1-3 are shown in Table 3 below.

Table 3
Tear strength (Die C)

Example No.	1	2	3
Carbon black, phr	80	88	95
Aromatic oil, phr	_ 22	0	22
Tear strength, N/mm	160	177	169

#### Comparative Examples 1-3

[0047] The elastomer composites of Comparative Examples 1-3 were prepared using a BR 1600 Banbury Mixer (Farrel). The mixing procedures are shown in Table 4.

Table 4
Dry mix preparation

Stage 1	Farrel BR Banbury Mixer (1600cc), 70% fill factor, 80 rpm, 45 °C
Time (min)	Operation
0	Add polymer
0.5	Add filler
2.5	Add oil (if any)
4	Sweep
5 or 8	Dump
	Pass through open mill 3 times
Stage 2	Farrel BR Banbury Mixer (1600 cc) 65% fill factor, 60 rpm, 45 °C.
Time (min)	Operation
0	Add Stage 1 compound and
	curatives
2	Dump
	Pass through open mill 3 times
	Sit at room temperature for at least 2 hours

[0048] The resulting elastomeric composites were tested for tear strength (Die C). The results are shown in Table 5.

Table 5
Tear strength (Die C)

Tour burongur (1510 C)					
Comparative Example No.	1	2	3		
Carbon black, phr Aromatic oil, phr	80 22	90 0	100 22		
Tear strength, N/mm	93	59	94		

[0049] As can be seen comparing the results shown in Table 3 with those in Table 5, elastomer composites of the present invention have considerably higher tear strength than those prepared using a conventional dry mixing method.

[0050] Although the present invention has been described above in terms of specific embodiments, it is anticipated that other uses, alterations and modifications thereof will become apparent to those skilled in the art given the benefit of this disclosure. It is intended that the following claims be read as covering such alterations and modifications as fall within the true spirit and scope of the invention.

#### [0051] What is claimed is:

1. An elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer, the particulate filler comprising at least one carbon black having structure and surface area values meeting the equation CDBP  $\leq$  (BET  $\div$  2.9).

- 2. An elastomer composite in accordance with claim 1 comprising at least 60 phr of the particulate filler.
- 3. An elastomer composite in accordance with claim 1 comprising at least 60 phr of the carbon black meeting the equation CDBP  $\leq$  (BET  $\div$  2.9).
- 4. An elastomer composite in accordance with claim 1 wherein the particulate filler further comprises at least one additional filler material having structure and surface area values not meeting the equation CDBP  $\leq$  (BET  $\div$  2.9).
- 5. An elastomer composite in accordance with claim 1 wherein the elastomer is selected from natural rubber, a homopolymer, copolymer or terpolymer of butadiene, styrene, isoprene, isobutylene, 2,3-dialkyl-1,3-butadiene where the alkyl group is C1 to C3 alkyl, acrylonitrile, ethylene or propylene.
- 6. An elastomer composite in accordance with claim 1 wherein the elastomer comprises natural rubber.
- 7. An elastomer composite in accordance with claim 1 further comprising at least one additive selected from antiozonants, antioxidants, plasticizers, processing aids, resins, flame retardants, extender oils, lubricants, and combinations thereof.
- 8. An elastomer composite in accordance with claim 1 wherein the elastomer composite has a tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm.

9. An elastomer composite comprising a general purpose rubber and particulate filler dispersed in the general purpose rubber, wherein the particulate filler comprises an amount of at least one carbon black effective in said general purpose rubber to achieve tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm.

10. An elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer, the elastomer composite having

Shore A hardness, measured in accordance with test method ASTM-D1415, greater than about 65;

tensile strength, measured in accordance with test method ASTM-D412, greater than 30 megapascals; and

elongation at break, measured in accordance with test method ASTM-D412, of at least about 600 %;

wherein the particulate filler comprises carbon black having structure and surface area values meeting the equation CDBP  $\leq$  (BET  $\div$  2.9).

- 11. An elastomer composite in accordance with claim 10 having tear strength, as measured by test method ASTM D-624, of at least 160 N/mm.
- 12. An elastomer composite comprising a general purpose rubber and particulate filler dispersed in the general purpose rubber, wherein the particulate filler comprises an amount of at least one carbon black effective in said general purpose rubber to achieve:

Shore A hardness, measured in accordance with test method ASTM-D1415, greater than about 65;

tensile strength, measured in accordance with test method ASTM-D412, greater than 30 megapascals; and

elongation at break, measured in accordance with test method ASTM-D412, of at least about 600 %.

13. An elastomer composite in accordance with claim 12 wherein said carbon black has structure and surface area values meeting the equation CDBP  $\leq$  (BET  $\div$  2.9).

- 14. An elastomer composite in accordance with claim 12 wherein said elastomer has tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm.
- 15. An elastomer composite in accordance with claim 1 wherein the particulate filler comprises at least one carbon black having structure and surface area values meeting the equation  $CDBP \le (BET \div 2.9) X$ , wherein X is about 5.
- 16. An elastomer composite in accordance with claim 1 wherein the particulate filler comprises at least one carbon black having structure and surface area values meeting the equation  $CDBP \le (BET \div 2.9) X$ , wherein X is about 10.
- 17. An elastomer blend comprising an elastomer composite blended with at least one elastomer material, the elastomer composite comprising a first elastomer and particulate filler dispersed in the first elastomer, the particulate filler comprising at least one carbon black having structure and surface area values meeting the equation  $CDBP \leq (BET \div 2.9)$ -X.
- 18. An elastomer blend in accordance with claim 17 wherein the elastomer material comprises elastomer different from the first elastomer.
- 19. An elastomer blend in accordance with claim 17 wherein the elastomer composite, before blending, has tear strength, as measured by test method ASTM-D624 using Die C, of at least about 160 N/mm.
- 20. An elastomer blend in accordance with claim 17 wherein the elastomer composite, before blending, has

Shore A hardness, measured in accordance with test method ASTM-D1415, greater than about 65;

tensile strength, measured in accordance with test method ASTM-D412, greater than 30 megapascals; and

elongation at break, measured in accordance with test method ASTM-D412, of at least about 600%.

21. A method of producing an elastomer composite comprising an elastomer and particulate filler dispersed in the elastomer, the method comprising:

feeding a continuous flow of first fluid comprising elastomer latex to a mixing zone; and

feeding a continuous flow of second fluid comprising the particulate filler under pressure to the mixing zone to form a mixture, the mixing of the first fluid and the second fluid within the mixing zone being sufficiently energetic to substantially completely coagulate the elastomer latex with the particulate filler;

the particulate filler comprising at least one carbon black having structure and surface area values meeting the equation CDBP  $\leq$  (BET  $\div$  2.9)-X.

- 22. The method of claim 21 wherein the elastomer composite has tear strength, as measured by test method ASTM-D624 using Die C, of at least 160 N/mm.
- 23. The method of claim 21 further comprising blending the elastomer composite with an elastomer material to form an elastomer blend.
- 24. The method of claim 23 wherein blending the elastomer composite with the elastomer material comprises dry mixing the elastomer composite with the elastomer material.
- 25. The method of claim 23 wherein the elastomer material comprises additional filler.

26. The method of claim 21 further comprising blending the elastomer composite with additional filler.

- 27. The method of claim 21 wherein the elastomer composite has tear strength, as measured by test method ASTM-D624 using Die C, of at least 160 N/mm.
- 28. The method of claim 21 wherein the elastomer composite has

  Shore A hardness, as measured by test method ASTM D1415, of
  at least 65,

tensile strength, as measured by test method ASTM D412, of at least 30 megapascals, and

elongation, as measured by test method ASTM D412, of at least 600 %.

- 29. An elastomer composite in accordance with claim 1, wherein the particulate filler comprises at least one carbon black having a CDBP value of about 43 to 45 mL/100g and a BET value of about 260 to 264 m<sup>2</sup>/g.
- 30. An elastomer composite in accordance with claim 29 wherein the at least one carbon black is present at a loading level from about 50 to 110 phr based upon the weight of the elastomer in the elastomer composite.

Intern: Application No PCT/US 02/37439

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C08L21/00 C08K3/00

C08K3/04

According to International Patent Classification (IPC) or to both national classification and IPC

#### B. FIELDS SEARCHED

 $\begin{array}{ccc} \text{Minimum documentation searched (classification system followed by classification symbols)} \\ \text{IPC 7} & \text{C08L} & \text{C08K} \end{array}$ 

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, PAJ

2010000-10	Carlos at de-		
Category °	Citation of document, with indication, where appropriate, of	the relevant passages	Relevant to claim No.
X	WO 00 34370 A (CABOT CORP) 15 June 2000 (2000-06-15)		1-8,10, 11, 15-20, 29,30
	page 6, line 1 -page 7, line	4	
X	US 6 042 643 A (GALLOWAY COLL 28 March 2000 (2000-03-28) cited in the application	IN P ET AL)	1-8,10, 11, 15-20, 29,30
	examples 47-59		
X	US 6 075 084 A (MABRY MELINDA 13 June 2000 (2000-06-13) cited in the application column 4, line 40 -column 5,		9,21-28
		-/	
X Furth	ner documents are listed in the continuation of box C.	χ Patent family members are	e listed in annex.
"A" docume consider the filling distribution of the filling distribution of the filling distribution of the filling docume of the fi	nt which may throw doubts on priority claim(s) or is cited to establish the publication date of another no rother special reason (as specified) ent referring to an oral disclosure, use, exhibition or	"T" later document published after it or priority date and not in conflicited to understand the principl invention  "X" document of particular relevance cannot be considered novel or involve an inventive step when  "Y" document of particular relevance cannot be considered to involve document is combined with onments, such combination being in the art.  "&" document member of the same	ict with the application but le or theory underlying the e; the claimed invention cannot be considered to the document is taken alone e; the claimed invention e an inventive step when the e or more other such docu- g obvious to a person skilled
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Interna ication No PCT/US 02/37439

<u> </u>		PCT/US 02	/3/439
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
X	US 5 100 637 A (NAKAI KIYONARI) 31 March 1992 (1992-03-31) example 4		12-14
A	US 5 830 930 A (FRANCIS ROBERT A ET AL) 3 November 1998 (1998-11-03) cited in the application  column 1, line 48 - line 51 table 2		1-8,10, 11, 15-20, 29,30
			*

nal application No. PCT/US 02/37439

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	ernational Searching Authority found multiple inventions in this international application, as follows:
	see additional sheet
1. X	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this international Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remar	k on Protest  The additional search fees were accompanied by the applicant's protest.  X  No protest accompanied the payment of additional search fees.
	X No protest accompanied the payment of additional search tees.

### FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-8, 10, 11, 15-30

Composite comprising (i) an elastomer and (ii) carbon black wherein the carbon black has a specific ratio of CDBP dibutylphthalate adsorption and BET surface area.

2. Claim: 9

Composite comprising (i) rubber and (ii) carbon black wherein the composite has a certain tear strength.

3. Claims: 12-14

Composite comprising (i) rubber and (ii) carbon black wherein the composite has a certain tensile strength, shore A hardness and elongation at break.

Intern: Application No
PCT/US 02/37439

Patent document		Publication		Patent family	02/3/439 Publication
cited in search report		date		member(s)	date
WO 0034370	Α	15-06-2000	AU	2354000 A	26-06-2000
			EP	1157063 A1	28-11-2001
			JP	2002531661 T	24-09-2002
		···	WO	0034370 A1	15-06-2000
US 6042643	Α	28-03-2000	US	5851280 A	22-12-1998
			US	6494946 B1	17-12-2002
			US AT	5900029 A 221106 T	04-05-1999 15-08-2002
			AU	706060 B2	10-06-1999
			AU	4378496 A	03-07-1996
			AU	706229 B2	10-06-1999
			AU	4472096 A	03-07-1996
			BR	9510016 A	28-10-1997
			CA	2207414 A1	20-06-1996
			CA	2207455 A1	20-06-1996
			CN	1321708 A	14-11-2001
			CN	1175272 A ,B	04-03-1998
			CN CZ	1175270 A ,B 9701841 A3	04-03-1998 12-11-1997
			CZ	9701841 A3 9701842 A3	15-07-1998
			DE	69527543 D1	29-08-2002
			DE	69527543 T2	03-04-2003
			DK	799281 T3	11-11-2002
			EG	20775 A	29-02-2000
			EG	21734 A	27-02-2002
			EP	1225205 A2	24-07-2002
			EP EP	0799281 A1 0797637 A1	08-10-1997 01-10-1997
			ES	2176357 T3	01-10-1997
			HU	76985 A2	28-01-1998
			JP	10510861 T	20-10-1998
			JP	10510863 T	20-10-1998
			NO	972736 A	13-08-1997
			NO	972737 A	08-08-1997
			NZ	298327 A	29-06-1999
			NZ PL	298988 A	28-10-1999
			PL	320731 A1 320830 A1	27-10-1997 10-11-1997
			RU	2157394 C2	10-11-1997
			SI	9520129 A	30-04-1998
			SI	9520133 A	30-06-1998
			TR	960556 A2	21-07-1996
			TR	960557 A2	21-07-1996
			TW	419510 B	21-01-2001
			WO WO	9618688 A1 9618696 A1	20-06-1996 20-06-1996
			US	5672198 A	20-06-1996 30-09-1997
			ZA	9510659 A	30-05-1996
			ZA	9510664 A	09-07-1996
US 6075084	Α	13-06-2000	US	6048923 A	11-04-2000
			US	5811642 A	22-09-1998
			AU	1063099 A	23-04-1999
			BR	9815397 A	11-12-2001
			CA	2305702 A1	08-04-1999
			CN EP	1280534 T 1019228 A1	17-01-2001 19-07-2000
			l- i	TOTATEO MI	13-0/-2000

Interna Application No
PCT/US 02/37439

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
US 6075084	A		JP	2001518401 T	16-10-2001
			PL	339614 A1	02-01-2001
			WO	9916600 A1	08-04-1999
			ÜS	2002016404 A1	07-02-2002
			ÜS	6413478 B1	02-07-2002
			AU	2600097 A	22-10-1997
			BR	9708412 A	24-10-2000
			CA	2250774 A1	09-10-1997
			EP	0892705 A2	27-01-1999
			JP	2000507892 T	27-06-2000
			KR	2000005190 A	25-01-2000
			SG	89248 A1	18-06-2002
			WO	9736724 A2	09-10-1997
			US	6040364 A	21-03-2000
			ZĀ	9702633 A	28-09-1998
					20 09 1990
US 5100637	Α	31-03-1992	JP	1913113 C	09-03-1995
			JP	2032137 A	01-02-1990
			JP	6041539 B	01-06-1994
			FR	2634492 A1	26-01-1990
			KR	9302555 B1	03-04-1993
US 5830930	Α	03-11-1998	ΑU	705605 B2	27-05-1999
			ΑU	6540396 A	11-12-1996
			BR	9609105 A	02-02-1999
			CA	2221564 Al	28-11-1996
			CN	1190977 A	19-08-1998
			CZ	9703682 A3	15-04-1998
			EP	0828789 A2	18-03-1998
			HU	9900397 A2	28-05-1999
			JP	3305723 B2	24-07-2002
			JP	11505879 T	25-05-1999
			NO	975336 A	16-01-1998
			NZ	313363 A	27-03-2000
			PL	323469 A1	30-03-1998
			SI	9620068 A	31-10-1998
			TR	9701405 T1	21-03-1998
			US	6008272 A	28-12-1999
			WO	9637547 A2	28-11-1996
			US	2003040553 A1	27-02-2003
			US	6323273 B1	27-11-2001
			US	6028137 A	22-02-2000
			US	2001036995 A1	01-11-2001
			US	5869550 A	09-02-1999
			US	5919841 A	06-07-1999
			ZA	9604060 A	27-08-1996